

Cosmopolitan Scolytinae: strong common drivers, but too many singularities for accurate prediction

Jean-Claude Grégoire¹, Hervé Jactel², Jiri Hulcr³, Andrea Battisti⁴,
Daegan Inward⁵, Françoise Petter⁶, Fabienne Grousset⁶

1 *Université libre de Bruxelles, 50 av. FD Roosevelt, 1050 Bruxelles, Belgium* **2** *INRAE, University of Bordeaux, umr Biogeco, Cestas, France* **3** *School of Forest, Fisheries and Geomatics Sciences, University of Florida, Gainesville, FL, USA* **4** *Università di Padova, DAFNAE-Entomologia, Agripolis 35020, Legnaro, Italia* **5** *Forest Research, Alice Holt Lodge, Farnham, Surrey, GU10 4LH, UK* **6** *European and Mediterranean Plant Protection Organization, Paris, France*

Corresponding author: Jean-Claude Grégoire (jean-claude.gregoire@ulb.be)

Academic editor: Marc Kenis | Received 1 July 2022 | Accepted 2 November 2022 | Published 18 May 2023

Citation: Grégoire J-C, Jactel H, Hulcr J, Battisti A, Inward D, Petter F, Grousset F (2023) Cosmopolitan Scolytinae: strong common drivers, but too many singularities for accurate prediction. In: Jactel H, Orazio C, Robinet C, Douma JC, Santini A, Battisti A, Branco M, Seehausen L, Kenis M (Eds) Conceptual and technical innovations to better manage invasions of alien pests and pathogens in forests. NeoBiota 84: 81–105. <https://doi.org/10.3897/neobiota.84.89826>

Abstract

Many scolytine beetle species have been expanding in new territories, travelling with wood and plants for planting, sometimes with a high impact on plant health. Here, we attempt to quantify the mobility of these species and to identify the biological drivers of mobility and impact. Mobility was estimated by counting the numbers of landmasses (contiguous pieces of land, surrounded by ocean or sea) colonised by each species. A series of potential drivers (taxonomic tribes; feeding regimes; polyphagy; reproductive strategy; host taxa; aggregation pheromones and long-range primary attractants), as well as impact on host health were recorded. A total of 163 species were identified, out of 5546 counted in the whole subfamily. The cosmopolitan taxa amongst the subfamily showed significant disharmony with regards to invasion frequency. Four tribes (Xyleborini; Ipini; Crypturgini; Hylastini) were significantly over-represented and two others (Corthylini; Hexacoloni) were under-represented. Some 53% of the 163 species are inbreeding, a very significant excess as compared to the whole subfamily (29%). The inbreeders colonised more landmasses than the outbreeders. There is a significant relationship between the number of host families attacked by a species and the number of colonised landmasses. Most of the invasive species are recorded to respond to long-range host primary attractants, only one quarter respond to pheromones. All very mobile species respond to long-range primary attractants and none is known to respond to pheromones. Very mobile species are all associated with a substantial or moderate impact. The most mobile species belong to a limited number of subtribes. They are often inbreeding, polyphagous and respond to long-range primary

attractants, but do not produce pheromones. However, there are many counter-examples. The outbreeding *Scolytus multistriatus* attacks only three host families, producing aggregation pheromones and has established in thirteen landmasses, with a high impact. Due to these many exceptions, species-based risk prediction relying on the few traits routinely analysed in literature suffers from important uncertainties.

Keywords

ambrosia beetles, bark beetles, entry, establishment, fungi, geographic distribution, impact, inbreeding, landmasses, polyphagy, quarantine, risk assessment, spread, semiochemicals

Introduction

Very few species are studied in depth before they become noticeable pests. Consequently, most attempts to assess the risk of potentially invasive species rely on limited information. Invasive species assessments now use multiple methodologies ranging from consensus-seeking horizon scans to climate match modelling. However, nearly all these methodologies suffer from one fundamental problem – lack of information about the interactions between a specific species and its potential new environment or hosts.

The typical solution is to take a broader taxonomic perspective and assume that the ecology of a species can be derived from the ecology of related species for which there is more knowledge or to assume that species within a genus are ecologically similar. The invasive species modelling literature is rich with examples of assessments of genera or even entire families or even guilds (see, for example, Mech et al. 2019; Barwell et al. 2020; Schulz et al. 2021).

Some of the most damaging forest pests in the world are bark and ambrosia beetles belonging to the weevil (Curculionidae) subfamily Scolytinae. Global climate change and intense silviculture enabled species, such as *Dendroctonus ponderosae* Hopkins and *Ips typographus* L., to multiply to epidemic proportions in North America and Europe, respectively (Grégoire et al. 2015; Hicke et al. 2016) with a total of 455.7 million m³ of pine killed by *D. ponderosae* in British Columbia between 2000 and 2015 (British Columbia Government 2019) and 148 million m³ of spruce killed by *I. typographus* between 1950 and 2000 in Europe (Schelhaas et al. 2003), with dramatically increasing damage during the last few years (Hlásny et al. 2021). In addition to these species which are currently spreading within their native continents only, many others have been expanding their territorial range worldwide, especially travelling with international trade. Bark and ambrosia beetles may additionally cause damage as vectors of pathogenic fungi. The redbay ambrosia beetle, *Xyleborus glabratus* Eichhoff, of Asian origin, was first reported in North America in 2002 (Rabaglia et al. 2006). It vectors the fungal symbiont, *Raffaelea lauricola* T.C. Harr., Fraedrich and Aghayeva, causing “laurel wilt”. At least 300 million *Persea borbonia* L. Spreng. (redbay trees) have been killed by laurel wilt in the USA (Hughes et al. 2017) and several other tree species of the Lauraceae family, including avocado (*Persea americana* Mill.) are also affected by the disease. Another example is the polyphagous shot hole borer, *Euwallacea fornicatus*

Eichhoff (Stouthamer et al. 2017; Smith et al. 2019), a Scolytinae which, together with a symbiotic *Fusarium* sp. fungus, attacks a large number of plants, mostly in cultivated settings in its area of origin, Asia, as well as, more recently, in areas of introduction¹ in North America (Rabaglia et al. 2006), Israel (Mendel et al. 2012) and South Africa (Paap et al. 2018). The North American species *Dendroctonus valens* LeConte was reported in China at the end of the 1990s and, by 2005, it had spread over 500,000 ha of pine forest in three provinces, killing more than 10 million *Pinus tabulaeformis* Carr. (Yan et al. 2005). Other harmful Scolytinae species killing living trees and recently introduced into Europe include the Asian ambrosia beetle, *Xylosandrus crassiusculus* (Motschulsky), the black twig borer, *X. compactus* (Eichhoff) and the walnut twig beetle, *Pityophthorus juglandis* Blackman. This latter species vectors the pathogenic fungus *Geosmithia morbida* Kolarík, Freeland, Utley and Tisserat, causing thousand cankers disease of walnuts, *Juglans* spp. (EPPO 2015; Seybold et al. 2019).

The observed spread of these species and many others continues. At the same time, dozens of bark- and ambrosia beetle species have been introduced into non-native regions without any detectable impact. Most bark beetle “tramp species” are harmless.

So far, at least 163 species out of the ~ 6,000 described scolytine species (Hulcr et al. 2015) are known to have established outside of their native areas (Table 1 and Suppl. material 1). The remaining ninety-eight percent of scolytine species are, thus, still potentially able to colonise new territories and their potential impact is still mostly unknown.

Other species that have not spread to date and which are not recognised as harmful, might start expanding their range, benefiting from the trade of new commodities or from commercial movements along new routes. These beetles, alone or together with pathogens, may also colonise new hosts that may prove to be more susceptible than their native hosts or form new associations with local pathogens as suggested by Rasati et al. (2019a). For both known and unrecognised spreading species, the possibility that they can be successfully introduced into new areas and their subsequent potential economic or environmental impact are two major components of phytosanitary risk.

“Horizontal” regulations globally addressing the host plants of non-native pests are locally implemented. For example, all non-European Scolytinae-attacking conifers are targeted in the European Union by phytosanitary requirements applying to the importation of coniferous wood² (EU 2019), but equivalent requirements do not exist for the trade of non-coniferous wood. A recent EPPO study focused on twenty-six representative Scolytinae and Platypodinae ambrosia- and bark beetle species associated with non-coniferous wood (EPPO 2020; Grousset et al. 2020). Sixteen life-history traits and other factors were qualitatively weighed with expert knowledge against invasion success. Inbreeding, polyphagy (number of host families) and the lack of aggregation pheromones were common features of species with a successful introduction

1 In this context, ‘the entry of a pest resulting in its establishment’, following the terminology of the Glossary of Phytosanitary Terms of the International Plant Protection Convention (FAO 2019).

2 ‘Commodities such as round wood, sawn wood, wood chips and wood residue, with or without bark, excluding wood packaging material, processed wood material and bamboo and rattan products’ (FAO 2021).

history. Association with pathogenic fungi, the use of aggregation pheromones and the capacity to attack and kill new host species were identified as factors contributing to high impact. One of the important conclusions of this EPPO study was that traits related to species with a past invasion history had a strong influence on invasion risks. However, it was found that the main factors that are driving successful establishment and impact vary from species to species and are not always fully identified. One important recommendation of this study was that horizontal phytosanitary measures similar to those for conifer wood better address the risk than regulation of individual species. In another recent study, Lantschner et al. (2020) similarly reviewed 123 Scolytinae species with a history of invasion, focusing on biological characteristics (feeding regime and mating strategy), cumulative trade between world regions, size of source species pools, forest area and climatic matching between the invaded and source regions. They identified sib-mating as a major factor favouring the movement of Scolytinae species into new territories, but also found that a non-biological trait, cumulative trade between world regions, is a primary driver of scolytine invasion.

At a broader taxonomic scale, Mech et al. (2019) and Schulz et al. (2021) focused on the impact of non-native herbivorous insects established in North America. They found that the evolutionary proximity between the native and novel host plants, life history traits of the novel hosts and the presence of native close congeners with a long-term association with the novel host were better predictors of impact than were traits of the invading insects themselves.

In this study, we tried to demonstrate that even relatively closely related species can differ in their capacity to colonise new territories and in their impact. We used presence in at least two landmasses (defined below) as a criterion to select 163 “mobile” Scolytinae species and quantified further their mobility by counting the number of colonised landmasses, according to the literature. We also ranked their impact on plant health. Finally, we attempted to identify biological and ecological features (feeding regimes, inbreeding, polyphagy, aggregation pheromones, primary attractants, conifer/non-conifer hosts or both), associated with differences in mobility and impact.

Methodology

The counting of colonised territories served as a proxy to estimate mobility. A dataset of Scolytinae species known to have spread beyond geographical barriers (across seas or oceans in this study) was constructed (Suppl. material 1), including any species distributed across at least one barrier (hereafter designated as “*Scolytinae with an invasion history*” - SIH), irrespective of its area of origin which is often difficult to delimit (see, for example, Lin et al. 2021). The list includes all the Scolytinae species from the EPPO study (EPPO 2020; Grousset et al. 2020), as well as the species introduced into North America, New Zealand and Europe, listed respectively by Haack (2001, 2006), Brockerhoff et al. (2006) and Kirkendall and Faccoli (2010). This initial set was expanded using information mostly from Wood and Bright (1992), Lantschner et al.

(2020), Atkinson (2021) and from other publications (full list of references in Suppl. material 1). The dataset was completed in December 2020 and, therefore, does not include several important studies (in particular Bright 2021) published after this date.

Amongst the biological features taken into account, the association with pathogens was not considered as a predictor because, in addition to previously known species, species so far harmless on their native hosts (e.g. *R. lauricola*, *G. morbida*) become pathogenic when their vectors colonise new host trees. Besides, scolytines species considered as harmless are sometimes found associated with aggressive pathogens (Wingfield and Gibbs 1991), making pathogens a dubious predictor of impact. Climatic requirements, dispersal capacity and voltinism were also not considered, because of the wide knowledge gap regarding these potential drivers (but see EPPO 2020 and Grousset et al. 2020).

Feeding regimes

We retained the following general categories (Kirkendall et al. 2015): phloeophagy (feeding in inner bark; this category corresponds to the bark beetles *stricto sensu*); xylomycetophagy (fungus farming; this category corresponds to the ambrosia beetles, which live in the xylem of woody plants, where they cultivate symbiotic fungi on which they feed); spermatophagy (feeding in seeds) and herbiphagy (feeding in non-woody plants).

Inbreeding

In some species, the females are fertilised by a brother, with extreme situations where the males are flightless and do not even leave their natal gallery. Only the species with full inbreeding were considered here. The outbreeding species that show some level of inbreeding (e.g. *Orthotomicus erosus*, *Tomicus piniperda*, *Hylurgus ligniperda*, *Ips grandicollis*, *Ips pini*) were not considered as inbreeding in this study. Unless specified in Suppl. material 1, the information comes from Kirkendall et al. (2015).

Polyphagy

Polyphagy was measured, as in EPPO (2020) and Grousset et al. (2020), by the number of host-plant families colonised. Unless specified otherwise in Suppl. material 1, host-plant data come from Wood and Bright (1992) or Atkinson (2021).

Aggregation pheromones (categories: 0/1/2)

We relied on published information, with the understanding that some species might use pheromones that have not been identified so far, for example, short-distance sex pheromones. The source for this field is El-Sayed (2018) unless specified otherwise. Three categories were considered: 0 (no pheromone identified or unknown for the genus); 1 (pheromones known for at least one other species in the genus); 2: (pheromone(s) identified in the species).

Long-range primary attractants (0/1/2)

Here too, we relied on published information, with the understanding that some species might respond to long-range primary attractants that have not been identified so far or only weakly respond to known attractants. Unless specified by a footnote in Suppl. material 1, the information regarding primary attractants (e.g. ethanol and/or alpha-pinene, emitted by the host or by other organisms within the host) comes from Atkinson (2021). The three categories considered are the same as for pheromones.

Host plants: conifers vs. non-conifers (1/2/3)

Three categories were considered: 1 (species attacking only conifers); 2 (species attacking only non-conifers); 3 (species attacking both conifers and non-conifers).

Impact on plant health (0/1/2)

Only direct impact on living trees and seeds or economic impact on traded products (e.g. logs, seeds) were considered. Other ecological impact mechanisms, such as those affecting native arthropods, wood decomposition and other ecosystem processes and patterns, can occur, but these are too poorly known in invasive Scolytinae. Even for the most obvious impact mechanisms, very few quantitative measurements are available in literature and only for a few species in a restricted number of areas. In addition, qualitative estimates vary greatly between assessors. We were, thus, led to rank impact according to three categories (0-1-2): 0 (no impact documented in the literature); 1 (moderate impact: some indication of impact, with some uncertainties because of discrepancies in literature); 2 (known substantial impact documented sometimes quantitatively by several sources). The criteria for damage by spermatophages were the reported colonisation of fruits (none – moderate – massive) and/or impact on regeneration (none – moderate – massive).

Landmasses

We use the term *landmass* to define a contiguous piece of land (a continent or an island, irrespective of its size) surrounded by ocean or sea. This approach admittedly creates large biases. Even if a continent is very large, we consider it as a single landmass. The movements of a species within a landmass are not considered because they are often incompletely documented. However, continents that are not fully separated by oceans (North, Central and South America; Europe, Asia and Africa) are considered as distinct landmasses because of the distances and ecoclimatic differences between them. Some archipelagos (e.g. Cape Verde, Fiji, Galápagos, Hawaii, Micronesia) were considered each as one unit. Islands comprising several countries (e.g. Republic of Ireland + Northern Ireland; Haiti + Dominican Republic) were considered as single units. The size of the geographic barriers between landmasses and of the landmasses themselves has not been considered. Great Britain and the European mainland would,

thus, be considered as separate landmasses, although the Channel that separates them is locally less than 35 km broad. On the other hand, South America, which is more than 7000 km long, is considered as a single landmass. Despite these many inconsistencies, we believe that this approach provides a useful, if probably conservative, metric to consider pest mobility. Suppl. material 1 provides a listing and a counting of the discrete landmasses occupied by each species. The acronyms used to designate the different landmasses are listed in Suppl. material 2. When possible, ISO alpha-3 codes (<https://www.iso.org/obp/ui/#search>) were used. Codes for locations absent from this list because they refer to intra-national territories (e.g. an island belonging to a larger country) were taken from the International Working Group on Taxonomic Databases For Plant Sciences (TDWG) (<https://github.com/tdwg/wgsrpd>) or were created for the purpose of this analysis.

Statistical analyses

Disharmony with regards to invasion frequency amongst the different SIH tribes; feeding regimes vs. reproductive strategies

2 × 2 Chi-Square tests were used, with Yate's correction for continuity for expected values inferior to 5.

Multivariate analyses on impact

A factorial discriminant analysis (FDA) was performed as a supervised classification method to discriminate amongst three categories of beetle species *a priori* classified, as in the Methodology and in Suppl. material 1, according to their level of damage (impact), as having no impact (0), moderate impact (1) or substantial impact (2), using ecological characteristics as predictor variables (Suppl. material 1). The dataset consisted of 163 species characterised by one quantitative functional trait, polyphagy, expressed as the number of known host plant families and five qualitative functional traits transformed into dummy variables, namely whether bark beetle species exhibited the following characteristics: xylomycetophagy (ambrosia beetles), inbreeding, using aggregation pheromones, using long-range primary attractants and host specialisation ("specialists": attacking either conifers or non-conifers; "generalists": attacking both) .

Covariance analyses on mobility

A Spearman correlation analysis was performed between the number of colonised land masses and the functional traits of the 163 scolytine species. Two variables were identified as significantly correlated with beetle cosmopolitanism, one quantitative, the degree of polyphagy (expressed in terms of number of known host plant families) and one qualitative, the use (or not) of long-range primary attractants for host plant colonisation. We then used an analysis of covariance (Ancova, with and without interaction) to assess the magnitude of the effects of these two factors. All statistical analyses were made with XLSTAT.

Results

Scolytinae with an invasion history - overall features

Disharmony with regards to invasion frequency amongst the SIH tribes

Five tribes, the Xyleborini, Trypophloeini, Ipini, Crypturgini and Hylastini are significantly more frequent amongst the invasive Scolytinae than amongst the Scolytinae as a whole. Two tribes, the Corthylini and Hexacolini are significantly less frequent (Table 1).

Tribes over-represented amongst the invasive Scolytinae are in bold, followed by (+); tribes under-represented are in bold, followed by (-). World figures taken from Hulcr et al. (2015), except for the Trypophloeini, Cryphalini, Corthylini and Ernoporini, for which the revision by Johnson et al. (2020a) was used. The number of non-SIH species is calculated by subtracting the number of SIH in a tribe from the total number of species in the tribe.

The small tribes Amphiscolytini (1 sp.), Cactopinini (21), Carphodicticini (5), Hyorrhynchini (19) and Phrixosomatini (25) are absent from the SIH list, as well as the larger tribes Diamerini (132), Micracidini (298) and Xyloctonini (78).

Feeding regimes

Amongst the 163 SIH species, 79 (48.5%) are phloeophagous, 60 (36.8%) are xy-lomycetophagous, twelve (7.4%) are herbiphagous and twelve are spermatophagous. The majority (82.3%) of the phloeophages amongst the SIH are outbreeding, whilst

Table 1. Tribes represented amongst the Scolytinae with an invasion history (SIH).

Tribes	SIH species		Non-SIH species		Total	Chi ²	
	N _{SIH}	Weight of tribe within category (%)	N _{non-SIH}	Weight of tribe within category (%)	N	Chi ² _(1, N)	p
Xyleborini (+)	56	34.4	1112	20.6	1168	17.0422	0.000024
Trypophloeini (+)	18	11.0	246	4.6	264	14.7696	0.000121
Dryocoetini	14	8.6	460	8.5	474	0.0004	0.984373
Ipini (+)	14	8.6	216	4.0	230	8.3351	0.003889
Crypturgini (+)	8	4.9	47	0.9	55	22.2837	< 0.00001
Scolytini	8	4.9	201	3.7	209	0.6013	0.438087
Hypoborini	7	4.3	202	3.7	209	0.1281	0.720387
Hylastini (+)	6	3.7	49	0.9	55	9.7088	0.001834
Hylurgini	6	3.7	124	2.3	130	0.7786	0.377564
Corthylini (-)	5	3.1	1237	22.9	1242	35.8508	< 0.00001
Cryphalini	5	3.1	247	4.6	252	0.8257	0.363514
Phloeosinini	4	2.5	223	4.1	227	1.1493	0.283696
Polygraphini	3	1.8	151	2.8	154	0.2465	0.619527
Hylesinini	2	1.2	162	3.0	164	1.1856	0.276221
Phloeotribini	2	1.2	108	2.0	110	0.1747	0.675995
Bothrosternini	1	0.6	130	2.4	131	1.5137	0.218568
Hexacolini (-)	1	0.6	241	4.5	242	5.6591	0.017365
Scolytplatypodini	1	0.6	52	1.0	53	0.0022	0.962393
Xyloterini	1	0.6	21	0.4	22	0.0344	0.852906
Ernoporini	1	0.6	177	3.3	178	2.8113	0.093603
Total	163	100	5406	100	5569		

Table 2. Feeding regimes of the Scolytinae with an invasion history.

Feeding regime	Outbreeding			Inbreeding			Total		Chi ² ₁	
	N	% of total	% of regime	N	% of total	% of regime	N	% of total	Chi ² ₁	p
Xylomycetophagy (+)	4	2.5	6.7	56	34.4	93.3	60	36.8	60.9222	< 0.00001
Phloeophagy (-)	65	39.9	82.3	14	8.6	17.7	79	48.5	78.3002	< 0.00001
Herbiphagy	5	3.1	41.7	7	4.3	58.3	12	7.4	0.128	0.720506
Spermatophagy (-)	2	1.2	16.7	10	6.1	83.3	12	7.4	4.6719	0.03066
Total	76	46.6		87	53.4		163			

the majority of the xylomycetophages (93.3%) and of the spermatophages (83.3%) are inbreeding. The mating habits of the herbiphages are equally balanced (Table 2).

Biological features influencing risks of introduction and impact

Mating strategy

Amongst the 163 species in our study, 87 (53.4%) are inbreeding (Table 3). This proportion of inbreeding species is significantly larger than that (27.8%) of the non-SIH inbreeders in the world (1544 species - Kirkendall et al. 2015) amongst the known species belonging to tribes with SIH species (5569 species - Hulcr et al. 2015; Johnson et al. 2020a): Chi²_(1; N=5569) = 47.42; p < 0.00001. The Xyleborini and Trypophloeini, over-represented in Table 1, are all inbreeding and the under-represented Corthylini and Hexacolini are all outbreeding. However, the over-represented Crypturgini and Hylastini are all outbreeding (Table 3).

Table 3. Mating strategies of the Scolytinae tribes with an invasion history.

Tribes	Outbreeding (% of tribe)	Inbreeding (% of tribe)	Total
Xyleborini	0	56	56
Trypophloeini	0	18	18
Cryphalini	5	0	5
Dryocoetini	4 (28.6%)	10 (71.4%)	14
Ipini	12 (85.7%)	2 (14.3%)	14
Crypturgini	8	0	8
Scolytini	8	0	8
Hypoborini	7	0	7
Hylastini	6	0	6
Hylurgini	5 (83.3%)	1 (16.7%)	6
Corthylini	5	0	5
Phloeosinini	4	0	4
Polygraphini	3	0	3
Hylesinini	2	0	2
Phloeotribini	2	0	2
Bothrosternini	1	0	1
Hexacolini	1	0	1
Scolytplatypodini	1	0	1
Xyloterini	1	0	1
Ernoporini	1	0	1
Total	76 (46.6%)	87 (53.4%)	163

Overall, the inbreeding (*stricto sensu*) SIH colonised a much larger set of landmasses than the outbreeding species (Fig. 1). Strikingly, with the exception of *Hypocryphalus mangiferae* (Stebbing) (17 landmasses), all the species colonising the larger numbers of landmasses are inbreeding.

Host nature and condition

The capacity to colonise living hosts appears to favour establishment. In our dataset, species with a recorded impact on their hosts colonised the larger numbers of landmasses (Fig. 2).

Host specificity

Amongst the 36 species in Suppl. material 1 attacking only conifers, 33 species attack only one family and two species attack two families. The Scolytinae attacking only non-conifers or attacking both non-conifers and conifers have a much wider and diverse range of host trees. Conifer specialists colonise fewer landmasses (median: 5) than non-conifer specialists (median: 6) and species attacking both types of hosts (median: 9) (see Fig. 3).

The genus *Hypothenemus*, representing 11% of the 163 species in the list, includes the most polyphagous species in the list with *H. eruditus*, reported from 65 plant families and *H. crudiae* and *H. seriatus*, each reported from 57 plant families. These species are reported from 37, 21 and 22 landmasses, respectively.

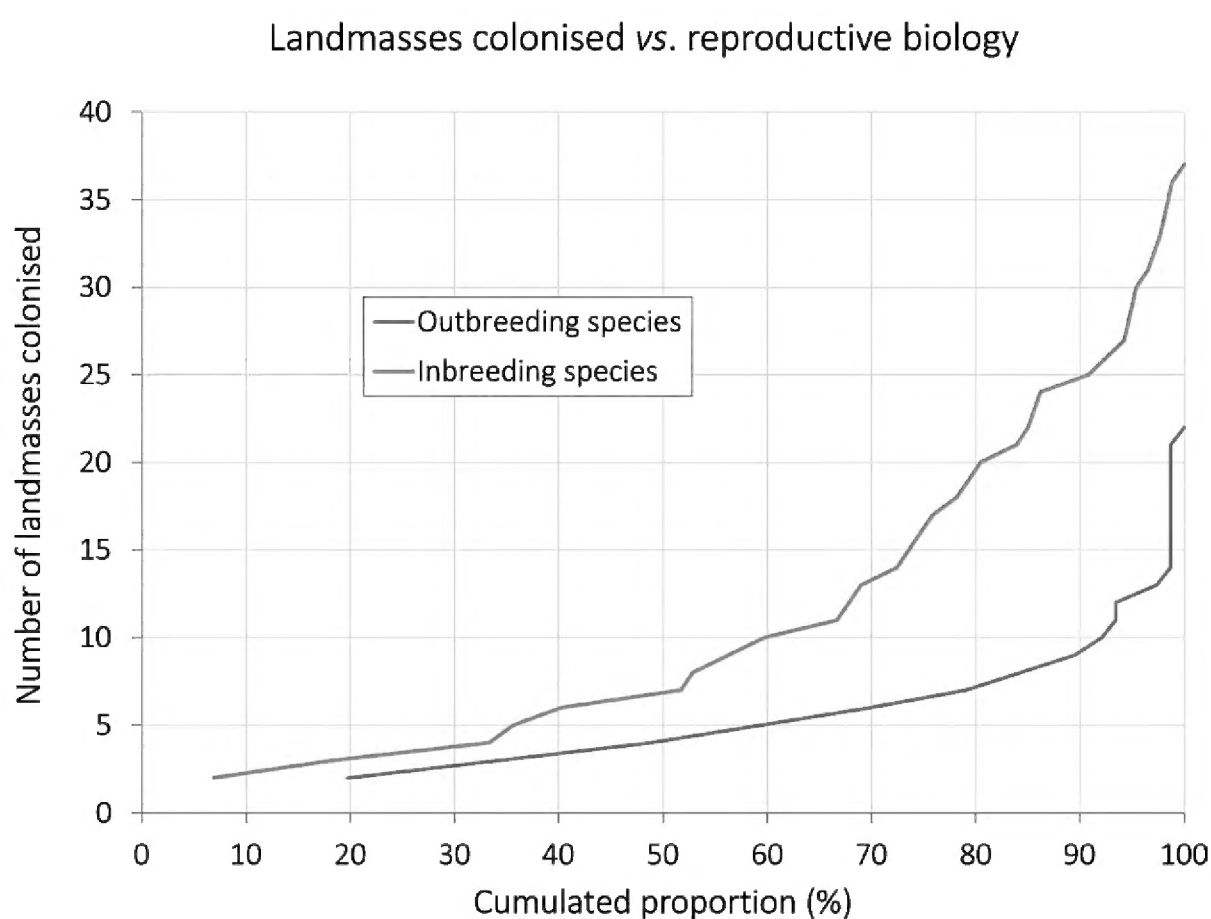


Figure 1. Cumulative proportion of landmasses colonised by either outbreeding or inbreeding species amongst the Scolytinae with an invasion history.

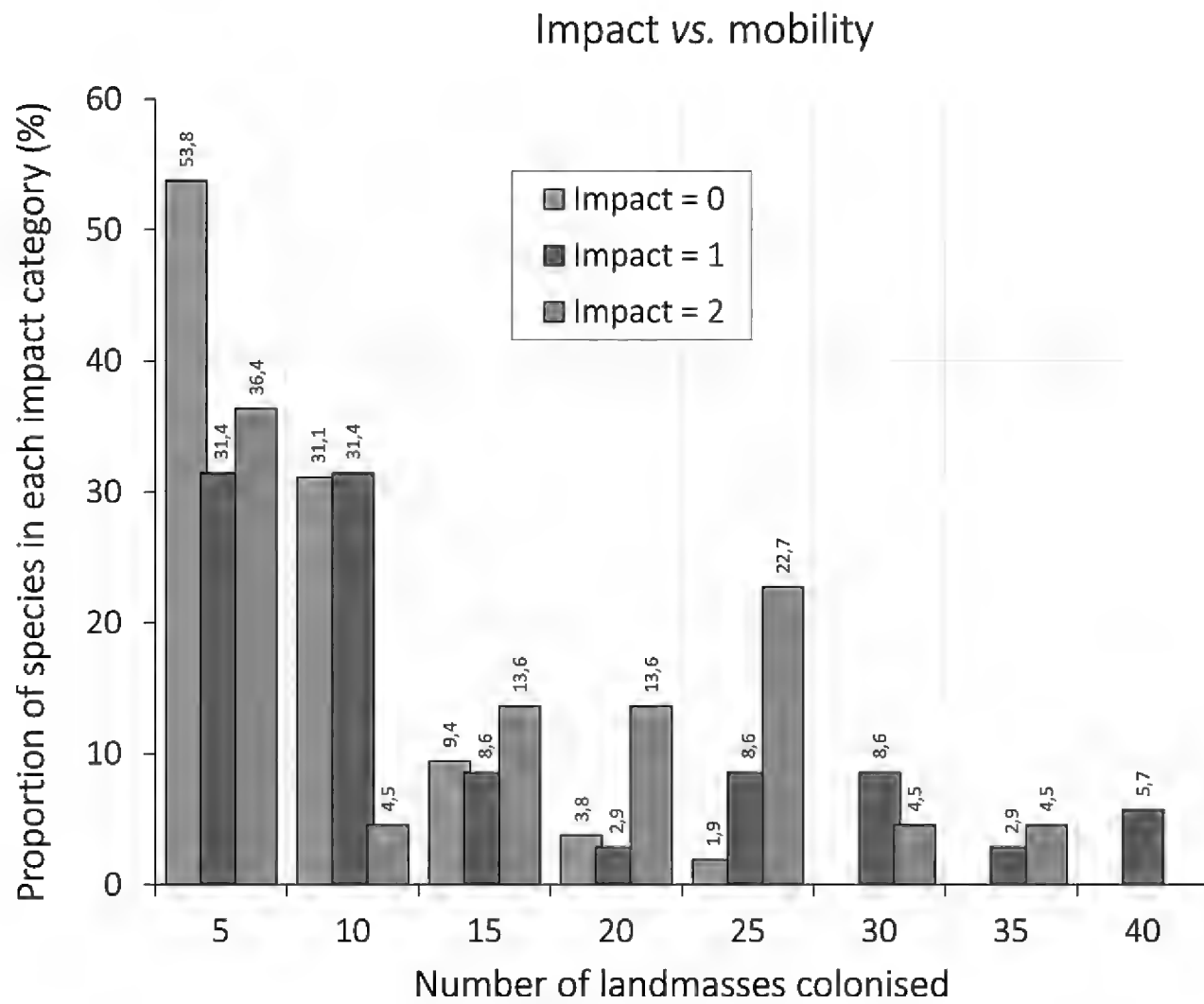


Figure 2. Impact versus mobility amongst scolytines with an invasion history.

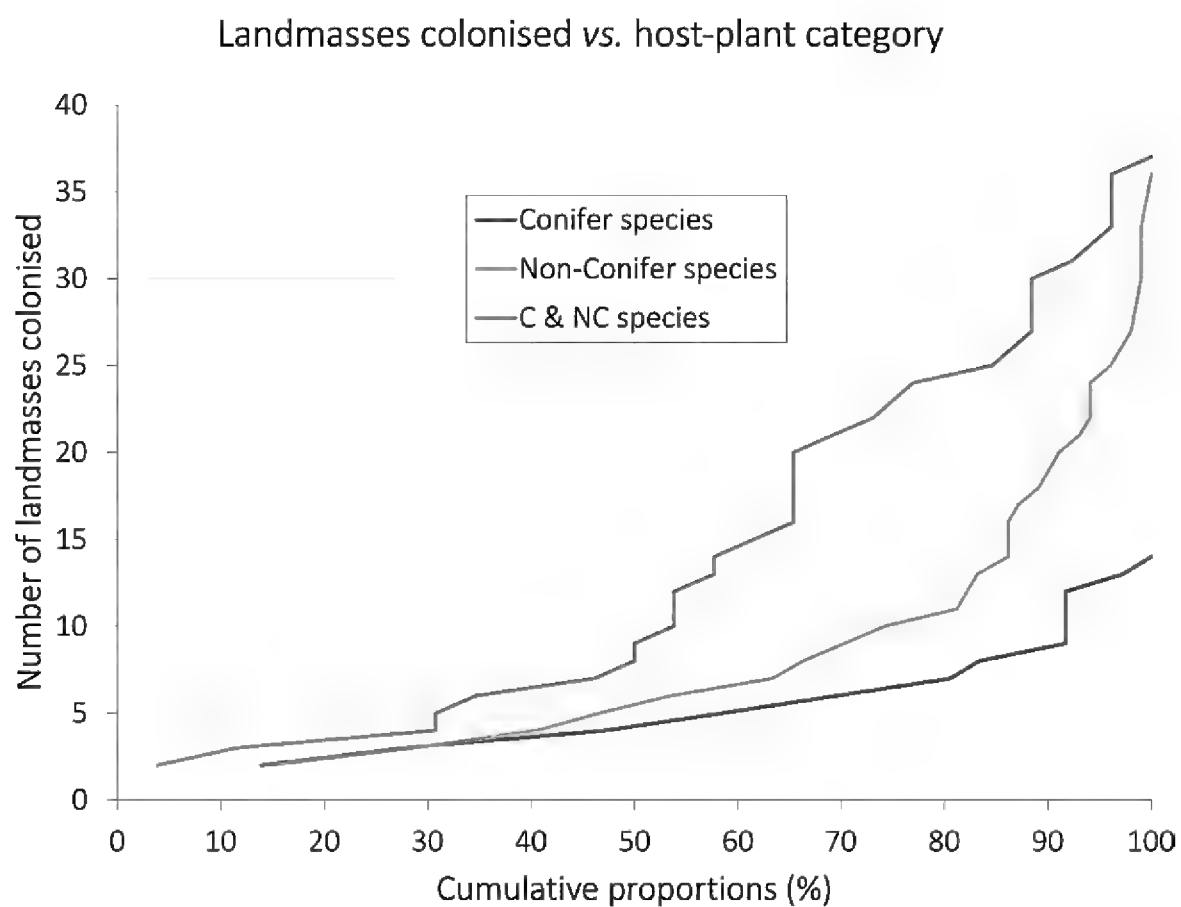


Figure 3. Host-plant category (conifer vs. non-conifer) influences the number of landmasses colonised by Scolytinae with an invasion history.

Aggregation pheromones and long-range primary attractants

Aggregation pheromones

Pheromone-mediated mass attacks are known amongst the SIH species, i.e. for *Orthotomicus erosus*, *Gnathotrichus materiarius* (Fitch), *Ips calligraphus* (Germar), *I. cembrae* (Herr), *I. grandicollis*, *Pityogenes bidentatus* (Herbst), *P. calcaratus* (Eichhoff), *P. chalcographus* (L.), *Pityokteines curvidens* (Germar), *Pityophthorus juglandis*, *Polygraphus poligraphus* (L.), *P. proximus* Blanford, *P. rufipennis* (Kirby), *Scolytus amygdali* Guerin-Meneville, *S. multistriatus* (Marsham), *T. domesticum* and many others.

Long-range primary attractants

94 SIH species out of 163 are known to respond to primary attractants and an additional 47 are likely to use these chemical clues as well.

Twenty species are not known to respond to primary attractants and do not produce pheromones either: five *Aphanarthrum* spp.; *Dendroctonus micans*; *Dryoxylon onoharaense*; *Kissophagus hederæ*; six *Liparthrum* spp.; *Microborus boops*; two *Microperus* spp; *Pagiocerus frontalis*; *Scolytoplatypus tycon*; *Thamnurgus characiae*.

Multivariate analyses

Impact

The factorial discriminant analysis showed significant effects of functional traits on impact (Wilks’ lambda test, $P < 0.0001$). The separation between the three impact levels was mainly explained by the FDA canonical function F1 (percentage variance explained 81.8%, $P < 0.0001$; while F2 explained 18.2%, $P = 0.09$). F1 was mainly driven by the degree of polyphagy ($P = 0.001$), use of aggregation pheromones ($P = 0.002$), host specialisation ($P = 0.004$) and, to a lesser extent, use of primary attractants ($P = 0.089$). The confusion matrix (Table 4) showed 100% correct classification for the category of non-damaging beetles (no impact; 107 species). The beetle species with no impact were characterised by a low degree of polyphagy, lack of aggregation pheromone, host specialisation on broadleaves or conifers and non-use of primary attractants. Only 11.4% of scolytine

Table 4. Confusion matrix for the factorial discriminant analysis (FDA) of the three categories of impact by the 163 beetle species studied.

<i>a priori</i> \ <i>a posteriori</i>	No impact	Low impact	Substantial impact	Total	% correct
No impact	107	0	0	107	100%
Moderate impact	29	4	2	35	11.4%
Substantial impact	15	4	2	21	9.5%
Total	151	8	4	163	69.3%

species with moderate impact and 9.5% with substantial impact were correctly classified, the other species of these categories being mainly misclassified as non-damaging. However, it should be noted that four *Euwallacea* species combined traits of polyphagy and lack of host specialisation, using aggregation pheromone and primary attractant: *E. piceus*, *E. interjectus*, *E. similis* and *E. validus* and they all had a significant impact.

The complete list of well-classified and misclassified species is available as supplementary material (Suppl. material 3).

Mobility

The Ancova analysis showed a significant effect of the degree of polyphagy ($P < 0.0001$) and use of primary attractant ($P = 0.023$) on the number of landmasses colonised, but the interaction of these two factors was not significant ($P = 0.58$), with an overall determination coefficient of $R^2 = 0.41$. Beetle species not using primary attractants ($n = 22$) colonised significantly fewer land masses (3.5 ± 0.4 , mean \pm standard error) than those ($n = 141$) attracted by the host plant (9.6 ± 0.7). The number of colonised landmasses increased with the degree of polyphagy (number of known host plant species) by the same magnitude for the two categories of beetle species (using or not primary attractants, Fig. 4).

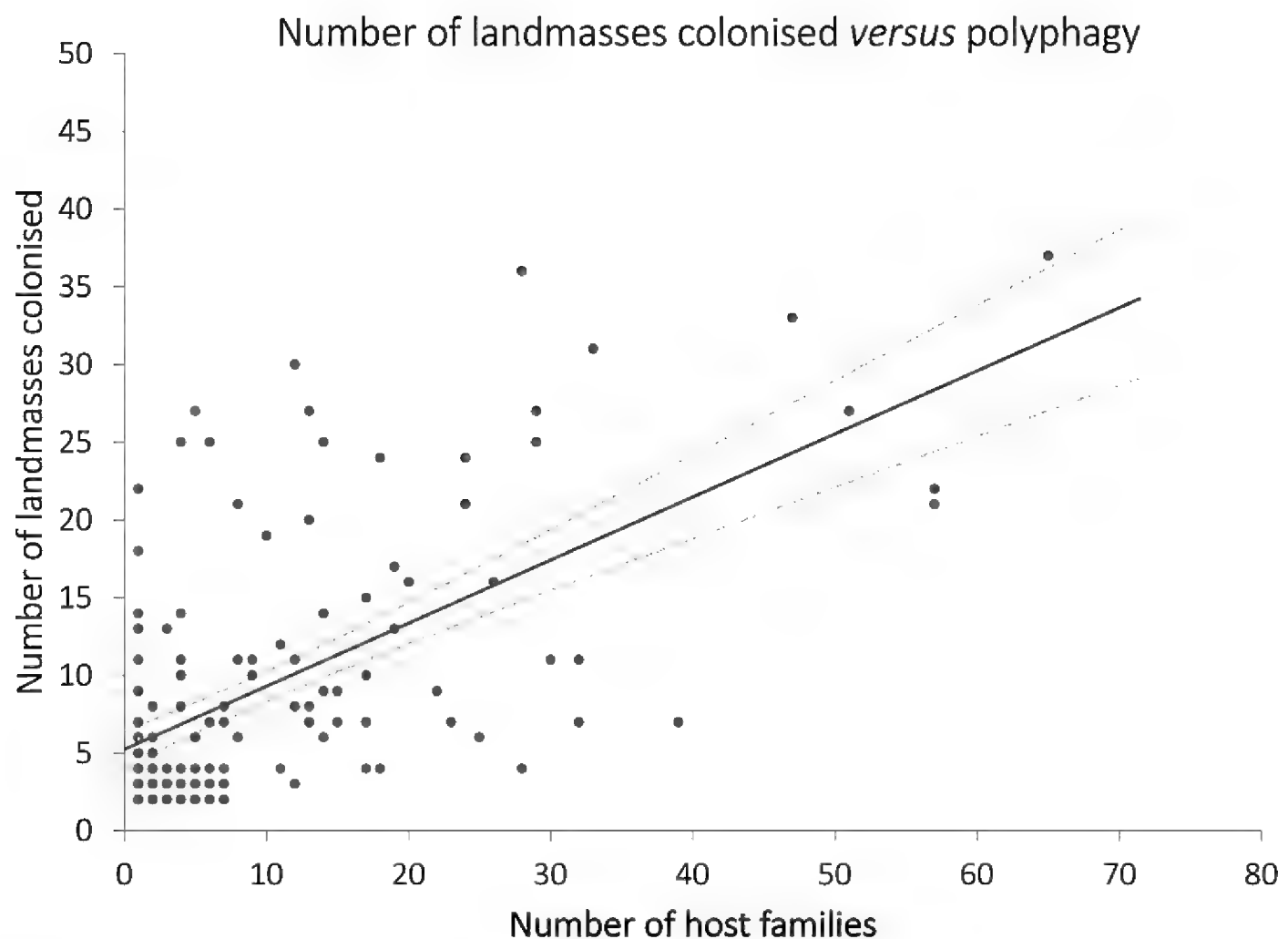


Figure 4. Number of colonised land masses versus degree of polyphagy (number of host-plant families) for the 163 scolytine species studied (independently of their use of primary attractants). Dashed lines represent the confidence interval of the linear regression line.

Discussion and conclusion

Disharmony with regards to invasion frequency amongst SIH tribes

Disharmony with regards to invasion frequency appears common amongst non-native insect orders worldwide and has been ascribed to the preference of certain orders for the main commercial pathways (Liebhold et al. 2016). Disharmony has been more finely recorded within Coleopteran (Liebhold et al. 2021) and Lepidopteran (Mally et al. 2022) non-native families.

Mating strategy

The SIH include a higher proportion of inbreeders than the world Scolytinae fauna. This is an asset for prompt establishment in strange lands. The females leaving the tree are already fertilised and can create a new colony on their own. In theory, the Allee population threshold (the minimal number of individuals below which a population cannot grow) for such species could be one single female.

Inbreeders are also often haplodiploid. Unfertilised females parthenogenetically produce haploid males and then mate with their sons (Jordal et al. 2000 and references therein). This further facilitates colonisation as females do not even have to be fertilised before dispersal and finding a host. For example, all the Xyleborini and most of the *Coccotrypes* spp. are haplodiploid (EPPO 2020; Grousset et al. 2020; Jordal et al. 2000).

Amongst the supposedly outbreeding species that crossed a geographic barrier, *Orthotomicus erosus* (Wollaston) (Mendel 1983) and *Tomicus piniperda* (Linnaeus) (Janin et al. 1988) show a proportion of females already mated upon emergence, possibly with a sibling or mated during maturation feeding on twigs or during overwintering at the base of trees previous to colonising a new host. Similarly, *Hylurgus ligniperda* (Fabricius) (Fabre and Carle 1975) and *Ips grandicollis* (Eichhoff) (Witanachchi 1980) have been observed to mate prior to emergence. As in the inbreeding species *stricto sensu*, these early mated females may be able to start a new colony alone. Wilkinson (1964) showed that *I. grandicollis* females induced alone to oviposit on pine logs produced a progeny. However, species with no invasive history are also capable of early mating. Lissimore (1997) found that three out of eight pre-emergent, overwintering *Ips pini* (Say) females collected in the spring in the litter around attacked trees were fertilised and able to start a new gallery alone. The North American species *Ips pini* has never expanded outside of its range, where it is widely distributed (Atkinson 2021). Similarly, Bleiker et al. (2013), examining 1510 emergent female *Dendroctonus ponderosae* Hopkins from two different locations in Alberta, found 3–5% of pre-emergent matings.

Host nature and condition

Many different relationships to the hosts are observed amongst bark- and ambrosia beetles, making it difficult to predict the risks associated with new insect-host associations or even the long-term risks associated with long-standing associations. Bark- and ambrosia

beetle species attack a wide range of trees, from apparently healthy individuals to dead and even decaying ones (Raffa et al. 2015; Hulcr et al. 2017). Other SIH species colonise a wide range of plant parts and, therefore, commodities in trade, including seeds, fine twigs and roots (Kirkendall et al. 2015 and see section 1.2). The nature and condition of the host allow a certain level of prediction regarding the entry, establishment and impact of a particular species or, after an event has occurred, provide clues for retrospective scenarios.

Entry

Xylophagous and xylomycetophagous species living in the sapwood are protected from mechanical damage and, when the wood has not been dried, from desiccation. Many phloeophagous bark beetles (e.g. *H. ligniperda*) and xylomycetophagous ambrosia beetles (e.g. *Xylosandrus germanus*) (Blandford) have travelled in wood packaging material or in wood or wood product shipments. The coffee berry borer, *Hypothenemus hampei* (Ferrari) is transported in the coffee seed trade (Johnson et al. 2020b). Plants for planting provide another pathways for species living in the stems of living hosts, such as *Xylosandrus compactus* (Eichhoff). *Coccotrypes dactyliperda* Fabricius, which live in dates, is likely to have spread around the world in commercial shipments. *C. rhizophorae* (Hopkins), which specifically lives in the propagules of the red mangrove, *Rhizophora mangle* Linnaeus, might have moved from Asia where it originates to North America in host propagules floating long distance across the ocean (Atkinson and Peck 1994).

Establishment and impact

Species capable of attacking living trees are more likely to find suitable hosts in the locations of entry. Hulcr et al. (2017) proposed to search for ambrosia beetle-fungus associations colonising live trees in their native habitats to identify future exotic tree-killing pests. Living trees, however, can vary in vigour and resistance to pests. Often, apparently healthy trees have been previously exposed to various forms of stress factors, including flooding, drought, wind break, snow break, freezing, ozone exposure, graft incompatibility, site and stand conditions, nutrients supply disorders, diseases or animal pest damage (Ranger et al. 2010; Ploetz et al. 2013; Hulcr and Stelinski 2017;) and this generally makes them more vulnerable to beetle attacks. Thirty-five SIH species may kill stressed hosts; twenty-one species out of 163 are able to kill apparently healthy, living trees (Suppl. material 1).

Importantly, the impact in a new area cannot always be predicted from the relationship of a beetle-fungus association with its native host trees. *X. glabratus* and its symbiont *R. lauricola* colonise stressed or injured Lauraceae all over the world. Whilst they exert little noticeable damage in their native areas, they massively kill *P. borbonia* in the USA because of the hypersensitive response of the New World Lauraceae and the changes in behaviour they induce in the beetles (Hulcr et al. 2017; Martini et al. 2017). *Anisandrus dispar* (Fabricius), which attacks weakened or dead trees in Europe is an important pest of young chestnut trees stressed by excess water or late frost in north-western USA and western Canada (Kühnholz et al. 2001). Similarly, *D. valens*, which usually settles on the stumps

of freshly cut pines or more rarely establishes in low numbers on stressed pines in North America, killed millions of *Pinus tabulaeformis* since its introduction into China during the late 1990s (Yan et al. 2005). The causes of this increased aggression in China are unclear, but have been related to exceptionally dry years following introduction (the outbreak subsided after the drought) and, possibly to some degree, to the association with a new, naïve host, with more aggressive strains of symbiotic fungi (Sun et al. 2013). Sometimes, even in their native range, species usually restricted to dead or dying hosts start attacking apparently healthy trees. *Trypodendron domesticum* (Linnaeus) and *T. signatum* (Fabricius) started infesting thousands of standing, live beech *Fagus sylvatica* L. in Belgium in the early 2000s, in connection with exceptional early frosts (La Spina et al. 2013). In Canada, *T. retusum* (LeConte) which is usually restricted to wind-broken or weakened trees was observed to attack apparently healthy aspen, *Populus tremuloides* Michaux (Kühnholz et al. 2001).

Scolytinae are not only a threat to forestry. For example, *H. hampei* is a major pest of coffee worldwide (Johnson et al. 2020b) and *C. dactyliperda* causes major damage on date production (Rodriguez et al. 2014).

Host specificity

Polyphagy and the ability to attack new hosts in new locations are advantageous for entry, establishment (higher probability of finding a suitable host) and impact (EPPO 2020).

Polyphagy

Bark beetles usually have a narrow host range and are often monophagous (all hosts belong to the same genus) or oligophagous (all hosts selected within one family). Ambrosia beetles often have a broader range of hosts, as their host is mainly a substrate for the fungi they grow and feed on (Beaver 1979; Jordal et al. 2000; Seybold et al. 2016). Many species specialise in either conifers or non-conifers, although some exceptionally polyphagous species attacks both.

There is no direct relationship between polyphagy and impact. Some less polyphagous ambrosia beetles have a substantial impact in newly-invaded territories, as illustrated by *X. glabratus* (4 host-plant families) after its introduction in the USA. On the contrary, very polyphagous species may cause limited damage in new areas, as well as in their native range. *Hypothenemus eruditus* (65 host families), which usually colonises dead hosts, is normally considered harmless (Kambestad et al. 2017).

New hosts

Many scolytines, even some not known as polyphagous, have been recorded on new host species when introduced into new areas (EPPO 2020; Grousset et al. 2020). Encounters with new hosts do not always result in damage, but are an important component of the potential impact. There are striking example of encounters with new very susceptible hosts, leading to extensive damage, such as *X. glabratus* on *Persea borbonia* in the USA (EPPO 2020) or *D. valens* on *P. tabulaeformis* in China (Yan et al. 2005).

Aggregation pheromones and long-range primary attractants

Aggregation pheromones

The need for mass-attacks can be unfavourable to establishment, but mass attacks, once the species is established and the epidemic threshold is reached, can result in higher impact (EPPO 2020). Some bark- or ambrosia beetles use aggregation pheromones to mass-attack standing hosts and overcome their defences (D.L. Wood 1982). The mass-colonisation of undefended, fallen trees is more likely the result of collective foraging, also mediated by aggregation pheromones (Toffin et al. 2018). As large numbers of individuals are required for a mass-attacking species to colonise a new tree, the Allee threshold is necessarily high, making establishment in a new area more difficult. On the contrary, solitary colonisers (e.g. *Hypothenemus* spp.; *Xylosandrus* spp.) have displayed high success in establishment (see section 2.1).

Long-range primary attractants

Physiologically stressed trees emit a range of volatile compounds, such as ethanol, which attract many bark- and ambrosia beetles colonising weakened hosts (Byers 1992; Miller and Rabaglia 2009; Ranger et al. 2010; Rassati et al. 2019b). Monoterpenes emitted by conifers also serve as clues for conifer-inhabiting species (Byers 1992), but reduce the response of species attacking non-conifers to ethanol or other primary attractants (Ranger et al. 2011). *H. hampei* is attracted to ripe coffee berries by conophthorin and chalcogran, but deterred by conifer monoterpenes (Jaramillo et al. 2013). Beetle response to primary attractants can be extremely accurate. In South Africa, Tribe (1992) showed that adults of the European species *Hylastes angustatus* (Herbst) and *Hylurgus ligniperda* were capable of finding *Pinus radiata* logs buried horizontally under 40 cm of soil. This accuracy is perhaps one component of the invasive success of these two species. However, working with native secondary species in Canada, Saint-Germain et al. (2007) showed that primary attractants allow bark beetles to locate a patch inhabited by susceptible hosts, but that, at closer range, host selection is governed by different processes, including random landing.

As they are not very specific (e.g. ethanol is produced by tissue fermentation of both conifers and non-conifers and monoterpenes, such as alpha-pinene, are produced by most conifers), long-range primary attractants can particularly facilitate host location and, thus, establishment amongst polyphagous species.

Conclusions

Throughout this review, several biological traits, particularly inbreeding and polyphagy, appear correlated with higher introduction potential and impact in new areas. However, as with the results obtained in EPPO (2020) and Grousset et al. (2020) for a narrower range of species, none of these traits, alone or combined, explains the success of all the SIH species and there are obvious outliers. For example, the over-represented

tribes Crypturgini and Hylastini (Table 1) are outbreeders. The moderately polyphagous *X. glabratus* (4 host families) has a much higher impact than *H. eruditus* (65 host families). More generally, 59 SIH species attack hosts in only one plant family, suggesting many exceptions to the influence of polyphagy on introduction. Whilst aggregation pheromones do not appear to favour establishment, there is the exception of *E. fornicatus*.

To summarise, some of the identified drivers are widespread amongst SIH species, but none is shared by the whole group, making it difficult to characterise univocally the potentially successful invaders amongst the bark- and ambrosia beetles of the world. In addition, the non-biological risk factors, as identified in EPPO (2020) and Lantschner et al. (2020), also play an important role. As concluded in EPPO (2020), the main factors that are driving successful establishment and impact vary from species to species and are not always fully identified. Still, one single feature common to most of the SIH species has been implicitly identified in this study on species crossing geographical barriers: their capacity to travel by trade, either on wood commodities and wood packaging material or on plants for planting or on fruits, depending on the species. The major conclusion of the present study is, thus, that, because of the lack of drivers that could allow for robust predictions regarding the invasive potential of any scolytine species, it is safer to consider the establishment of horizontal measures for trade of commodities.

Acknowledgements

HJ and AB participated in this study as part of the HOMED project (<http://homed-project.eu/>), which received funding from the European Union's Horizon 2020 Research and Innovation Programme, under grant agreement no. 771271. We are grateful to Sandy Liebhold and Miloš Knížek for useful comments on an earlier draft.

References

- Atkinson TH (2021) Bark and Ambrosia Beetles. <http://www.barkbeetles.info/index.php> [accessed on 10 February 2021]
- Atkinson TH, Peck SB (1994) Annotated checklist of the bark and ambrosia beetles (Coleoptera: Platypodidae and Scolytidae) of tropical southern Florida. *The Florida Entomologist* 77(3): 313–329. <https://doi.org/10.2307/3496101>
- Barwell LJ, Perez-Sierra A, Henricot B, Harris A, Burgess TI, Hardy G, Scott P, Williams N, Cooke DEL, Green S, Chapman DS, Purse BV (2020) Evolutionary trait-based approaches for predicting future global impacts of plant pathogens in the genus *Phytophthora*. *Journal of Applied Ecology*: 1–13. <https://doi-org.ezproxy.ulb.ac.be/10.1111/1365-2664.13820>
- Beaver RA (1979) Host specificity of temperate and tropical animals. *Nature* 281(5727): 139–141. <https://doi.org/10.1038/281139a0>
- Bleiker KP, Heron RJ, Braithwaite EC, Smith GD (2013) Preemergence mating in the mass-attacking bark beetle, *Dendroctonus ponderosae* (Coleoptera: Curculionidae). *Canadian Entomologist* 145(1): 12–19. <https://doi.org/10.4039/tce.2012.102>

- Bright DE (2021) Catalog of Scolytidae (Coleoptera), supplement 4 (2011–2019) with an annotated checklist of the world fauna (Coleoptera: Curculionoidea: Scolytidae). Doctoral dissertation, Colorado State University Libraries. <https://mountainscholar.org/handle/10217/229307>
- British Columbia Government (2019) MPB projections. <https://www2.gov.bc.ca/gov/content/industry/forestry/managing-our-forest-resources/forest-health/forest-pests/bark-beetles/mountain-pine-beetle/mpb-projections?keyword=mountainandkeyword=pineandkeyword=beetleandkeyword=2016> [accessed on 21 July 2019]
- Brockhoff EG, Bain J, Kimberley M, Knížek M (2006) Interception frequency of exotic bark and ambrosia beetles (Coleoptera: Scolytinae) and relationship with establishment in New Zealand and worldwide. *Canadian Journal of Forest Research* 36(2): 289–298. <https://doi.org/10.1139/x05-250>
- Byers JA (1992) Attraction of bark beetles, *Tomicus piniperda*, *Hylurgops palliatus*, and *Trypodendron domesticum* and other insects to short-chain alcohols and monoterpenes. *Journal of Chemical Ecology* 18(12): 2385–2402. <https://doi.org/10.1007/BF00984957>
- El-Sayed AM (2018) The Pherobase: Database of Pheromones and Semiochemicals. <http://www.pherobase.com> [consulted on 29 November 2020]
- EPPO (2015) Mini data sheet on *Geosmithia morbida* and *Pityophthorus juglandis*. https://gd.eppo.int/download/doc/955_minids_GEOHMO_en.pdf
- EPPO (2020) EPPO Study on the Risk of Bark and Ambrosia Beetles Associated with Imported Non-Coniferous Wood. EPPO Technical Document No. 1081; EPPO: Paris, France. https://www.eppo.int/RESOURCES/eppo_publications [accessed on 20 November 2020]
- EU (2019) Commission Implementing Regulation (EU) 2019/2072 of 28 November 2019 establishing uniform conditions for the implementation of Regulation (EU) 2016/2031 of the European Parliament and the Council, as regards protective measures against pests of plants, and repealing Commission Regulation (EC) No 690/2008 and amending Commission Implementing Regulation (EU) 2018/2019.
- Fabre JP, Carle P (1975) Contribution à l'étude biologique d'*Hylurgus ligniperda* F. (Coleoptera Scolytidae) dans le Sud-est de la France. *Annales des Sciences Forestières* 32(1): 55–71. <https://doi.org/10.1051/forest/19750104>
- FAO (2021) ISPM 5 Glossary of phytosanitary terms. IPPC, FAO, Rome.
- Grégoire J-C, Raffa KF, Lindgren BS (2015) Economics and Politics of Bark Beetles. In: Vega F, Hofstetter R (Eds) *Bark Beetles, Biology and Ecology of Native and Invasive Species*, 1st edn., Academic Press, 585–613. <https://doi.org/10.1016/B978-0-12-417156-5.00015-0>
- Grousset F, Grégoire J-C, Jactel H, Battisti A, Benko Beloglavec A, Hrašovec B, Hulcr J, Inward D, Orlinski A, Petter F (2020) The Risk of Bark and Ambrosia Beetles Associated with Imported Non-Coniferous Wood and Potential Horizontal Phytosanitary Measures. *Forests* 11(3): 342. <https://doi.org/10.3390/f11030342>
- Haack RA (2001) Intercepted Scolytidae (Coleoptera) at U.S. ports of entry: 1985–2000. *Integrated Pest Management Reviews* 6(3/4): 253–282. <https://doi.org/10.1023/A:1025715200538>
- Haack RA (2006) Exotic bark-and wood-boring Coleoptera in the United States: Recent establishments and interceptions. *Canadian Journal of Forest Research* 36(2): 269–288. <https://doi.org/10.1139/x05-249>

- Hicke JA, Meddens AJ, Kolden CA (2016) Recent tree mortality in the western United States from bark beetles and forest fires. *Forest Science* 62(2): 141–153. <https://doi.org/10.5849/forsci.15-086>
- Hlásny T, König L, Krokene P, Lindner M, Montagné-Huck C, Müller J, Qin H, Raffa KF, Schelhaas MJ, Svoboda M, Viiri H, Seidl R (2021) Bark beetle outbreaks in Europe: State of knowledge and ways forward for management. *Current Forestry Reports* 7(3): 138–165. <https://doi.org/10.1007/s40725-021-00142-x>
- Hughes MA, Riggins JJ, Koch FH, Cognato AI, Anderson C, Formby JP, Dreaden T, Ploetz R, Smith JA (2017) No rest for the laurels: Symbiotic invaders cause unprecedented damage to southern USA forests. *Biological Invasions* 19(7): 2143–2157. <https://doi.org/10.1007/s10530-017-1427-z>
- Hulcr J, Stelinski LL (2017) The Ambrosia Symbiosis: From Evolutionary Ecology to Practical Management. *Annual Review of Entomology* 62(1): 285–303. <https://doi.org/10.1146/annurev-ento-031616-035105>
- Hulcr J, Atkinson TH, Cognato AI, Jordal BH, McKenna DD (2015) Morphology, taxonomy, and phylogenetics of bark beetles. In: Vega F, Hofstetter R (Eds) *Bark Beetles, Biology and Ecology of Native and Invasive Species*, 1st edn., Academic Press, 41–84. <https://doi.org/10.1016/B978-0-12-417156-5.00002-2>
- Hulcr J, Black A, Prior K, Chen CY, Li HF (2017) Studies of ambrosia beetles (Coleoptera: Curculionidae) in their native ranges help predict invasion impact. *The Florida Entomologist* 100(2): 257–261. <https://doi.org/10.1653/024.100.0219>
- Janin JL, Lieutier F, Garcia J (1988) Existence de fécondations précoces dans le cycle biologique de *Tomicus piniperda* L. (Coleoptera Scolytidae) en forêt d'Orléans (France). *Agronomie* 8(2): 169–172. <https://doi.org/10.1051/agro:19880211>
- Jaramillo J, Torto B, Mwenda D, Troeger A, Borgemeister C, Poehling HM, Francke W (2013) Coffee berry borer joins bark beetles in coffee klatch. *PLoS ONE* 8(9): e74277. <https://doi.org/10.1371/journal.pone.0074277>
- Johnson AJ, Hulcr J, Knížek M, Atkinson TH, Mandelshtam MY, Smith SM, Jordal BH (2020a) Revision of the bark beetle genera within the former Cryphalini (Curculionidae: Scolytinae). *Insect Systematics and Diversity* 4(3): 1–81. <https://doi.org/10.1093/isd/ixaa002>
- Johnson MA, Ruiz-Diaz CP, Manoukis NC, Verle Rodrigues JC (2020b) Coffee berry borer (*Hypothenemus hampei*), a global pest of coffee: Perspectives from historical and recent invasions, and future priorities. *Insects* 11(12): 882. <https://doi.org/10.3390/insects11120882>
- Jordal BH, Normark BB, Farrell BD (2000) Evolutionary radiation of an inbreeding haplodiploid beetle lineage (Curculionidae, Scolytinae). *Biological Journal of the Linnean Society* 71(3): 483–499. <https://doi.org/10.1111/j.1095-8312.2000.tb01270.x>
- Kambestad M, Kirkendall LR, Knutsen IL, Jordal BH (2017) Cryptic and pseudo-cryptic diversity in the world's most common bark beetle – *Hypothenemus eruditus*. *Organisms, Diversity & Evolution* 17(3): 633–652. <https://doi.org/10.1007/s13127-017-0334-6>
- Kirkendall LR, Faccoli M (2010) Bark beetles and pinhole borers (Curculionidae, Scolytinae, Platypodinae) alien to Europe. *ZooKeys* 56: 227–251. <https://doi.org/10.3897/zookeys.56.529>
- Kirkendall LR, Biedermann PHW, Jordal BH (2015) Evolution and Diversity of Bark and Ambrosia Beetles. In: Vega F and Hofstetter R (Eds) *Bark Beetles, Biology and Ecology of*

- Native and Invasive Species, 1st edn., Academic Press, 85–156. <https://doi.org/10.1016/B978-0-12-417156-5.00003-4>
- Kühnholz S, Borden JH, Uzunovic A (2001) Secondary ambrosia beetles in apparently healthy trees: Adaptations, potential causes and suggested research. *Integrated Pest Management Review* 6(3/4): 209–219. <https://doi.org/10.1023/A:1025702930580>
- La Spina S, De Cannière C, Dekri A, Grégoire J-C (2013) Frost increases beech susceptibility to scolytine ambrosia beetles. *Agricultural and Forest Entomology* 15(2): 157–167. <https://doi.org/10.1111/j.1461-9563.2012.00596.x>
- Lantschner MV, Corley JC, Liebhold AM (2020) Drivers of global Scolytinae invasion patterns. *Ecological Applications* 30(5): e02103. <https://doi.org/10.1002/eap.2103>
- Liebhold AM, Yamanaka T, Roques A, Augustin S, Chown SL, Brockerhoff EG, Pyšek P (2016) Global compositional variation among native and non-native regional insect assemblages emphasizes the importance of pathways. *Biological Invasions* 18(4): 893–905. <https://doi.org/10.1007/s10530-016-1079-4>
- Liebhold AM, Turner RM, Blake RE, Bertelsmeier C, Brockerhoff EG, Nahrung HF, Pureswaran DS, Roques A, Seebens H, Yamanaka T (2021) Invasion disharmony in the global biogeography of native and non-native beetle species. *Diversity and Distributions* 27(11): 2050–2062. <https://doi.org/10.1111/ddi.13381>
- Lin W, Park S, Jiang ZR, Ji Y, Ernstsons AS, Li J, Li Y, Hulcr J (2021) Native or Invasive? The Red-Haired Pine Bark Beetle *Hylurgus ligniperda* (Fabricius)(Curculionidae: Scolytinae) in East Asia. *Forests* 12(7): 950. <https://doi.org/10.3390/f12070950>
- Lissemore FM (1997) Frass clearing by male pine engraver beetles (*Ips pini*; Scolytidae): Paternal care or paternity assurance? *Behavioral Ecology* 8(3): 318–325. <https://doi.org/10.1093/beheco/8.3.318>
- Mally R, Turner RM, Blake RE, Fenn-Moltu G, Bertelsmeier C, Brockerhoff EG, Hoare RJB, Nahrung HF, Roques A, Pureswaran DS, Yamanaka T, Liebhold AM (2022) Moths and butterflies on alien shores: Global biogeography of non-native Lepidoptera. *Journal of Biogeography* 49(8): 1455–1468. <https://doi.org/10.1111/jbi.14393>
- Martini X, Hughes MA, Killiny N, George J, Lapointe SL, Smith JA, Stelinski LL (2017) The fungus *Raffaelea lauricola* modifies behavior of its symbiont and vector, the redbay ambrosia beetle (*Xyleborus glabratus*), by altering host plant volatile production. *Journal of Chemical Ecology* 43(5): 519–531. <https://doi.org/10.1007/s10886-017-0843-y>
- Mech AM, Thomas KA, Marsico TD, Herms DA, Allen CR, Ayres MP, Gandhi KJK, Gurevitch J, Havill NP, Hufbauer RA, Liebhold AM, Raffa KF, Schultz AN, Uden DR, Tobin PC (2019) Evolutionary history predicts high-impact invasions by herbivorous insects. *Ecology and Evolution* 9(21): 12216–12230. <https://doi.org/10.1002/ece3.5709>
- Mendel Z (1983) Seasonal history of *Orthotomicus erosus* (Coleoptera: Scolytidae) in Israel. *Phytoparasitica* 11(1): 13–24. <https://doi.org/10.1007/BF02980707>
- Mendel Z, Protasov A, Sharon M, Zveibil A, Ben Yehuda S, O'Donnell K, Rabaglia R, Wysoki M, Freeman S (2012) An Asian ambrosia beetle *Euwallacea fornicatus* and its novel symbiotic fungus *Fusarium* sp. pose a serious threat to the Israeli avocado industry. *Phytoparasitica* 40(3): 235–238. <https://doi.org/10.1007/s12600-012-0223-7>

- Miller DR, Rabaglia RJ (2009) Ethanol and (-)- α -pinene: Attractant kairomones for bark and ambrosia beetles in the southeastern US. *Journal of Chemical Ecology* 35(4): 435–448. <https://doi.org/10.1007/s10886-009-9613-9>
- Paap T, De Beer ZW, Migliorini D, Nel WJ, Wingfield MJ (2018) The polyphagous shot hole borer (PSHB) and its fungal symbiont *Fusarium euwallaceae*: A new invasion in South Africa. *Australasian Plant Pathology* 47(2): 231–237. <https://doi.org/10.1007/s13313-018-0545-0>
- Ploetz RC, Hulcr J, Wingfield MJ, de Beer ZW (2013) Destructive Tree Diseases Associated with Ambrosia and Bark Beetles: Black Swan Events in Tree Pathology? *Plant Disease* 97(7): 856–872. <https://doi.org/10.1094/PDIS-01-13-0056-FE>
- Rabaglia RJ, Dole SA, Cognato AI (2006) Review of American Xyleborina (Coleoptera: Curculionidae: Scolytinae) occurring north of Mexico, an illustrated key. *Annals of the Entomological Society of America* 99(6): 1034–1056. [https://doi.org/10.1603/0013-8746\(2006\)99\[1034:ROAXCC\]2.0.CO;2](https://doi.org/10.1603/0013-8746(2006)99[1034:ROAXCC]2.0.CO;2)
- Raffa KF, Grégoire J-C, Lindgren BS (2015) Natural History and Ecology of Bark Beetles. In: Vega F, Hofstetter R (Eds) *Bark Beetles, Biology and Ecology of Native and Invasive Species*, 1st edn., Academic Press, 1–40. <https://doi.org/10.1016/B978-0-12-417156-5.00001-0>
- Ranger CM, Reding ME, Persad AB, Herms DA (2010) Ability of stress-related volatiles to attract and induce attacks by *Xylosandrus germanus* (Coleoptera: Curculionidae, Scolytinae) and other ambrosia beetles. *Agricultural and Forest Entomology* 12: 177–185. <https://doi.org/10.1111/j.1461-9563.2009.00469.x>
- Ranger CM, Reding ME, Gandhi KJ, Oliver JB, Schultz PB, Canas L, Herms DA (2011) Species dependent influence of (-)- α -pinene on attraction of ambrosia beetles (Coleoptera: Curculionidae: Scolytinae) to ethanol-baited traps in nursery agroecosystems. *Journal of Economic Entomology* 104(2): 574–579. <https://doi.org/10.1603/EC10243>
- Rassati D, Marini L, Malacrino A (2019a) Acquisition of fungi from the environment modifies ambrosia beetle mycobiome during invasion. *PeerJ* 7: e8103. <https://doi.org/10.7717/peerj.8103>
- Rassati D, Marini L, Marchioro M, Rapuzzi P, Magnani G, Poloni R, Di Giovanni F, Mayo P, Sweeney J (2019b) Developing trapping protocols for wood-boring beetles associated with broadleaf trees. *Journal of Pest Science* 92(1): 267–279. <https://doi.org/10.1007/s10340-018-0984-y>
- Rodriguez M, Delibes M, Fedriani JM (2014) Hierarchical levels of seed predation variation by introduced beetles on an endemic Mediterranean palm. *PLoS ONE* 9(10): e109867. <https://doi.org/10.1371/journal.pone.0109867>
- Saint-Germain M, Buddle CM, Drapeau P (2007) Primary attraction and random landing in host-selection by wood-feeding insects: A matter of scale? *Agricultural and Forest Entomology* 9(3): 227–235. <https://doi.org/10.1111/j.1461-9563.2007.00337.x>
- Schelhaas MJ, Nabuurs GJ, Schuck A (2003) Natural disturbances in the European forests in the 19th and 20th centuries. *Global Change Biology* 9(11): 1620–1633. <https://doi.org/10.1046/j.1365-2486.2003.00684.x>
- Schulz AN, Mech AM, Ayres MP, Gandhi KJ, Havill NP, Herms DA, Hoover AM, Hufbauer RA, Liebhold AM, Marsico TD, Raffa KF, Tobin PC, Uden DR, Thomas KA (2021) Predicting non-native insect impact: Focusing on the trees to see the forest. *Biological Invasions* 23(12): 3921–3936. <https://doi.org/10.1007/s10530-021-02621-5>

- Seybold SJ, Penrose RL, Graves AD (2016) Invasive bark and ambrosia beetles in California Mediterranean forest ecosystems. Insects and diseases of Mediterranean forest systems. Springer International Publishing, Switzerland, 583–662. https://doi.org/10.1007/978-3-319-24744-1_21
- Seybold S, Klingeman W III, Hishinuma S, Coleman T, Graves A (2019) Status and Impact of Walnut Twig Beetle in Urban Forest, Orchard, and Native Forest Ecosystems. *Journal of Forestry* 117(2): 152–163. <https://doi.org/10.1093/jofore/fvy081>
- Smith SM, Gomez DE, Beaver RA, Hulcr J, Cognato AI (2019) Reassessment of the Species in the *Euwallacea fornicatus* (Coleoptera: Curculionidae: Scolytinae) Complex after the Rediscovery of the “Lost” Type Specimen. *Insects* 10(9): 261. <https://doi.org/10.3390/insects10090261>
- Stouthamer R, Rugman-Jones P, Thu PQ, Eskalen A, Thibault T, Hulcr J, Wang L-J, Jordal BJ, Chen C-Y, Cooperband M, Lin CS, Kamata N, Lu S-S, Masuya H, Mendel Z, Rabaglia R, Sanguansub S, Shih H-H, Sittichaya W, Zong S (2017) Tracing the origin of a cryptic invader: phylogeography of the *Euwallacea fornicatus* (Coleoptera: Curculionidae: Scolytinae) species complex. *Agricultural and Forest Entomology* 19(4): 366–375. <https://doi.org/10.1111/afe.12215>
- Sun J, Lu M, Gillette NE, Wingfield MJ (2013) Red turpentine beetle: Innocuous native becomes invasive tree killer in China. *Annual Review of Entomology* 58(1): 293–311. <https://doi.org/10.1146/annurev-ento-120811-153624>
- Toffin E, Gabriel E, Louis M, Deneubourg J-L, Grégoire J-C (2018) Colonization of weakened trees by mass-attacking bark beetles: No penalty for pioneers, scattered initial distributions and final regular patterns. *Royal Society Open Science* 5(1): 170454. <https://doi.org/10.1098/rsos.170454>
- Tribe GD (1992) Colonisation sites on *Pinus radiata* logs of the bark beetles, *Orthotomicus erosus*, *Hylastes angustatus* and *Hylurgus ligniperda* (Coleoptera: Scolytidae). *Journal of the Entomological Society of Southern Africa* 55(1): 77–84. https://hdl.handle.net/10520/AJA00128789_3199
- Wilkinson C (1964) Attraction and development of *Ips* bark beetles in artificially infested pine bolts exposed on fire towers and turntables in Florida. *The Florida Entomologist* 47(1): 57–64. <https://doi.org/10.2307/3493769>
- Wingfield MJ, Gibbs JN (1991) *Leptographium* and *Graphium* species associated with pine infesting bark beetles in England. *Mycological Research* 95(11): 1257–1260. [https://doi.org/10.1016/S0953-7562\(09\)80570-4](https://doi.org/10.1016/S0953-7562(09)80570-4)
- Witanachchi JP (1980) Evidence for pre-emergence mating among mature progeny of *Ips grandicollis* (Eichhoff). *Australian Journal of Entomology* 19(2): 93–100. <https://doi.org/10.1111/j.1440-6055.1980.tb00969.x>
- Wood DL (1982) The role of pheromones, kairomones, and allomones in the host selection and colonization behavior of bark beetles. *Annual Review of Entomology* 27(1): 411–446. <https://doi.org/10.1146/annurev.en.27.010182.002211>
- Wood SL, Bright Jr DE (1992) A catalog of Scolytidae and Platypodidae (Coleoptera), Part 2. Taxonomic Index (Volumes A, B). *Great Basin Naturalist Memoirs* 13: 1–1553. <https://biostor.org/reference/143868>
- Yan Z, Sun J, Owen D, Zhang Z (2005) The red turpentine beetle, *Dendroctonus valens* LeConte (Scolytidae): An exotic invasive pest of pine in China. *Biodiversity and Conservation* 14(7): 1735–1760. <https://doi.org/10.1007/s10531-004-0697-9>

Supplementary material 1

Main characteristics of the 163 Scolytinae with an invasion history

Authors: Jean-Claude Grégoire, Hervé Jactel, Jiri Hulcr, Andrea Battisti, Daegan Inward, Françoise Petter, Fabienne Grousset

Data type: table (docx file)

Explanation note: Main characteristics of the 163 Scolytinae with an invasion history.

Copyright notice: This dataset is made available under the Open Database License (<http://opendatacommons.org/licenses/odbl/1.0/>). The Open Database License (ODbL) is a license agreement intended to allow users to freely share, modify, and use this Dataset while maintaining this same freedom for others, provided that the original source and author(s) are credited.

Link: <https://doi.org/10.3897/neobiota.84.89826.suppl1>

Supplementary material 2

Landmasses

Authors: Jean-Claude Grégoire, Hervé Jactel, Jiri Hulcr, Andrea Battisti, Daegan Inward, Françoise Petter, Fabienne Grousset

Data type: table (docx file)

Explanation note: Landmasses (islands and continents).

Copyright notice: This dataset is made available under the Open Database License (<http://opendatacommons.org/licenses/odbl/1.0/>). The Open Database License (ODbL) is a license agreement intended to allow users to freely share, modify, and use this Dataset while maintaining this same freedom for others, provided that the original source and author(s) are credited.

Link: <https://doi.org/10.3897/neobiota.84.89826.suppl2>

Supplementary material 3

Well-classified and misclassified species

Authors: Jean-Claude Grégoire, Hervé Jactel, Jiri Hulcr, Andrea Battisti, Daegan Inward, Françoise Petter, Fabienne Grousset

Data type: table (docx file)

Explanation note: Well-classified and misclassified species identified by the factorial discriminant analysis (FDA) of the three categories of impact by the 163 beetle species studied.

Copyright notice: This dataset is made available under the Open Database License (<http://opendatacommons.org/licenses/odbl/1.0/>). The Open Database License (ODbL) is a license agreement intended to allow users to freely share, modify, and use this Dataset while maintaining this same freedom for others, provided that the original source and author(s) are credited.

Link: <https://doi.org/10.3897/neobiota.84.89826.suppl3>